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- Catalytic process for preparing cyclohexanone-oxime.
- A catalytic process for preparing cyclohexanoneoxime by reacting cyclohexanone with NH₂ and H₂O₂ in the liquid phase, wherein the catalyst substantially consists of a highly crystalline substance containing SiO₂ and having a zeolitic structure.

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CATALYTIC PROCESS FOR PREPARING CYCLOHEXANONE-OXIME

DE-C-1 245 371 teaches the preparation of cyclohexanoneoxime by catalytic reaction, in the liquid phase, of cyclohexanone with ammonia and hydrogen peroxide at 5 to 40°C, in the presence of a catalyst consisting of phospho-tungstic acid or of similar compounds. A drawback of this method is, however, that this type of catalyst is difficult to handle, particularly, during the separation of the product from the catalyst.

It has now been found that it is possible to obtain the same high yields by using a completely different catalyst that is easier to handle and simplifies the process in all its steps.

Thus, the present invention relates to a catalytic process for preparing cyclohexanone-oxime by reacting cyclohexanone with NH₂ and H₂O₂ in the liquid phase, which process is characterized in that the catalyst substantially consists of a highly crystalline, SiO₂-containing substance having a zeolitic structure and, in particular, of a titanium-silicalite, optionally in admixture with an inert binder. Titanium-silicalites are known compounds which, for example, are described in GB-A-2 024 790 and 2 071 071.

The reaction temperature normally is in the range of from 25 to 100°C, preferably from 40 to 90°C. Tests carried out at 15°C have provided results that are not completely satisfactory.

A pressure above atmospheric pressure promotes the development of the reaction.

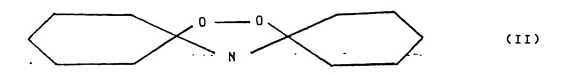
Other substances of zeolitic nature which catalyse the reaction are, first of all, the various types of silicalite, such as silicalite I (see, e.g., US-A-4 061 724), silicalite II, zirconium-silicalites and hafnium-silicalites.

Another class of analogous catalysts are the metalsilicalites, e.g., borosilicates (boralites), beryllosilicates, chromo-silicates, vanadium-silicates, zirconiumsilicates, gallium-silicates and ferro-silicates which are, in part, described in GB-A-2 024 790.

A third class of catalysts of analogous type consists of the known aluminium-silicates, generally known as "zeolites", particularly the zeolites of type Y, the zeolites ZSM5, the zeolites ZSM 11 and the other zeolites ZSM described in EP-A-129 239, 141 514 and 143 642, as well as the zeolites MB 28 described in EP-A-21 445. Some of these zeolites have already been successfully utilized in the aminoximation of cyclohexanone in the gas phase with NH3 and air.

The process according to the present invention can be carried out either continuously or discontinuously, provided that reactors with surfaces that are resistant to hydrogen peroxide are used. When the reaction is carried out in batch, it is advisable that 0.1 to 50 parts by weight (preferably from 1 to 20 parts by weight) of pure catalyst (excluding binder) for 100 parts by weight of cyclohexanone be used. If the reaction is performed in a continuous manner, it is preferred to employ a space velocity of from 0.1 to 100 kg/h of cyclohexanone -(C₆H₁₀O) per kg of catalyst. The molar ratio of H₂O₂:C₆H₁₀O is, generally, in the range of from 0.5 to 2.5 and, preferably, from 1 to 1.5, whereby H₂O₂ means 100% pure hydrogen peroxide (i.e., dilution water excluded). Water (H2O) is the most suitable liquid vehicle for the reaction. However, it is also possible to use organic water-soluble solvents capable of dissolving both pure ammonia and its aqueous solutions such as, e.g., methanol, ethanol, propanol, isopropanol, butanol, isobutanol, tertiary butanol and mixtures thereof. Solvents with carbonyl functions such as aldehydes and ketones should be excluded.

The reaction water, which is formed according to the equation: $C_{\epsilon}H_{10}O + H_{2}O_{2} + NH_{3} \rightarrow C_{\epsilon}H_{10} = NOH + 2H_{2}O$ (I) gradually increases the amount of liquid vehicle as the conversion to oxime proceeds. During the reaction considerable amounts of peroxy-di-cyclohexyl-amine of the formula



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are also formed.

Ammonia should always be added before the hydrogen peroxide and in excess (NH₂: C₄H₁₀O ≥1, preferably ≥ 1.5 moles/mol). Otherwise, undesired side-reactions occur.

At the end of the reaction, the cyclohexanoneoxime can be separated by different methods, for example, through extraction with suitable solvents such as benzene, toluene and the cyclohexanone as used for the synthesis, whereby a hydrophobic organic phase and an aqueous phase are formed.

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Cyclohexanone-oxime and unreacted cyclohexanon form the organic layer. The aqueous layer which contains the excess NH₃ as well as traces of cycloh xanone and oxime can be recycled to the reaction zone (ammoximation zone).

The following examples illustrate the invention without, however, limiting it.

EXAMPLE 1

A glass reactor, equipped with a stirrer and a heating jacket, was first blanketed with an inert gas (nitrogen). Subsequently, 1.5 g of a catalyst consisting of a finely ground powder of titaniumsilicalite containing 3.85% by weight of highly crystalline titanium oxide (average diameter of the particles ≤5 µm was charged. 50 cm³ of a 32% by weight aqueous solution of ammonia were added to the catalyst. The mixture was stirred and 9.5 g of cyclohexanone were charged. The three-phase system (solid-aqueous-organic) formed was kept homogenous by stirring vigorously. Heating was started by conveying a liquid at 60°C into the jacket. Simultaneously, a 32% by weight aqueous solution of hydrogen peroxide was fed to the reactor by means of a metering pump. After 15 minutes, the temperature in the reactor reached 60°C while the pressure rose to a value of from 600 to 700 mm Hg above atmospheric pressure. The addition of H2O2 was performed within 3.5 hours. during which time the pressure decreased. The temperature was maintained at 60°C and stirring was continued for a further 1.5 hours, whereafter the test was stopped and the mixture was cooled.

50 cm³ of toluene were added to the resulting suspension and, after stirring for a few minutes, the aqueous phase and the organic phase were separated from the catalyst by filtration. The liquid phases were separated in a separation funnel and

the aqueous phase was extracted with two portions of toluene of 30 cm³ each. The toluene solutions were combined and analysed. The analysis revealed a cyclohexanone conversion of 95% and a selectivity to cyclohexanone-oxime of 79.45%; the selectivity of hydrogen peroxide to oxime was 68.7%. At the end, about 15% of the starting cyclohexanone was found to be in the form of peroxy-dicyclohexyl-amine (see formula II above).

EXAMPLES 2 and 3

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Example 1 was repeated, varying temperature and pressure. The data and results given in table 1 prove that the selectivity of the conversion $H_2O_2 \rightarrow$ oxime is very adversely affected when operating under vacuum.

EXAMPLE 4 (Comparative Test)

Example 1 was repeated, bringing the pressure (gauge) to zero and considerably lowering the temperature (down to 15°C) The unsatisfactory results shown in table 1 prove that it is disadvantageous to excessively reduce the thermal level of the ammoximation.

EXAMPLE 5

Example 1 was repeated, replacing titanium silicalite by a zirconite (zirconium-silicalite). Analogous results were obtained.

EXAMPLE 6

Example 1 was repeated, replacing titaniumsilicalite by a boralite (boron-silicate). Analogous results were obtained.

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TABLE 1 (a)

	1	PARAMETERS	ETER	S			1		PRODUCTS	TS			2 6 6 8 8 8
	C H C	H,0,	1	Time	م	Liquid	iquid phase	Gas composition	osition	Conversi	on (%)	Conversion (%) Selectivity (%)	Lty (%)
Ëx.	01.9	6 10 2 2 (°C) (h)	ິ)	(h)	(mmHg)	(mmHg) aqueous	organic	N20	² 0	UHJ	UH	UHJ	O H
	ĥ		•		(p)	(a)	(g)	(vol.%)	(vol.%)	(vol.%) 6"10" "2"2	1,5,5	_e102_2	1,5,5
	9,50	9,50 11,7	09	3 1/2	7.5	52.0 127	127	17,22	23,14	95,35	91,87	79,45	68,73
2		9,50 11,55	40	3	negat	50.8	50,8 114,5	ı		90,37	79,20	30,97	25,74
					176								
. ო	9,50	11.7	52	=	0	57.0	120	0,335	95,49	82,56	98,13	20,53	15,37
4	9,55	11.8 15	12	3 3/4	0	71,3	16	0.094	20.69	47.79	95,48	16,70	7.22
*					1	1	1			1	1		

(*) Comparative test.

In all the tests, 50 cm 3 of aqueous NII $_3$ were added and 1.5 g of Ti-silicalite were used. (a)

(b) Above atmospheric pressure.

(c) Selectivity to oxime.

Note : Nitrogen protoxide derives from the decomposition of hydroxylamine, which forms in consequence of a parallel reaction.

Claims

- 1. A catalyst process for preparing cycloh xanone-oxim by reacting cyclohexanon with ammonia and hydrogen peroxide in the liquid phase, characterized in that the catalyst substantially consists of a highly crystalline substance containing SiO_z and having a zeolitic structure.
- 2. The process of claim 1, wherein the catalyst is a titanium-silicalite, optionally in admixture with an inert binder.
- 3. The process of claim 2, wherein a zirconium-silicalite replaces, at least in part, titanium-silicalite.
- 4. The process of claim 2, wherein a boralite replaces, at least in part, titanium-silicalite.

- 5. The process of any one of claims 1 to 4, wherein the reaction temperature ranges from 25 to 100°C.
- 6. The process of claim 5, wherein the reaction temperature ranges from 40 to 90°C.
- 7. The process of any one of claims 1 to 6, wherein the pressure is equal to or higher than atmospheric pressure.
- 8. The process of any one of claims 1 to 7, wherein the space velocity of cyclohexanone ranges from 0.1 to 100 kg/h per kg of pure catalyst (binder excluded).
- 9. The process of any one of claims 1 to 8, wherein the H₂O₂: C₄H₁₀O molar ratio ranges from 0.5 to 2.5, whereby H₂O₂ means pure hydrogen peroxide (dilution water excluded).
- 10. The process of any one of claims 1 to 9, wherein NH₃ is added prior to H_2O_2 , the NH₃: $C_4H_{10}O$ molar ratio being ≥ 1 , preferably ≥ 1.5 .

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